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VIRULENCE-ASSOCIATED ADHESINS

All documents cited herein are incorporated by reference in their entirety.

TECHNICAL FIELD

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This invention is in the field of bacterial adhesion. In particular, it relates to virulence-related adhesion antigens derived from *Haemophilus influenzae*, *Escherichia coli* and other organisms.

BACKGROUND ART

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The Gram negative Haemophilus genus includes H.influenzae, H.aegyptius (also referred to as H.influenzae biogroup aegyptius), H.decreyi and H.somnus. These bacteria can cause diseases including conjunctivitis, chancroid, purpuric fever, meningitis, pneumonia and epiglottitis. H.influenzae is the most commonly-found pathogen in this genus, and includes both typeable (encapsulated) and non-typeable (non-capsulated; 'NTHi') strains.

A vaccine against *H.influenzae* type B ('Hib') based on a conjugate of its capsular saccharide and a carrier protein has been enormously successful, but there has been little progress in providing protection against other members of the species. In particular, type D *H.influenzae* and non-typeable *H.influenzae* remain problematic.

Similarly, vaccines remain unavailable for other bacterial pathogens such as enterotoxigenic (ETEC), enteropathogenic (EPEC), enteroaggregative (EAEC), enterohemorrhagic (EHEC) and shiga-toxic (STEC) strains of *Escherichia coli*.

It is an object of the invention to provide materials and methods to improve the prevention and treatment of infections caused by such bacteria. More particularly, it is an object of the invention to provide materials suitable for immunising against bacterial infections.

DISCLOSURE OF THE INVENTION

Virulence-associated antigens involved in adhesion have been identified in several bacteria and other organisms, and these antigens are useful for the diagnosis, prevention and treatment of bacterial infections (particularly those caused by virulent strains). In particular, antigens have been identified in: Haemophilus influenzae biogroup aegyptius (SEQ ID NO: 1); Escherichia coli K1 (SEQ ID NO^S: 2 & 3) and also in EHEC strain EDL933; Actinobacillus actinomycetemcomitans (SEQ ID NO: 4); Haemophilus somnus (SEQ ID NO: 5); Haemophilus ducreyi (SEQ ID NO: 6); EPEC E.coli strain E2348/69 (SEQ ID NO: 7); EPEC (SEQ ID NO: 18); EAEC E.coli strain O42 (SEQ ID NO^S: 8 & 9); uropathogenic E.coli (SEQ ID NO: 10); Shigella flexneri (SEQ ID NO: 11); Brucella melitensis (SEQ ID NO: 12); Brucella suis (SEQ ID NO: 13); Ralstonia solanacearum (SEQ ID NO: 14); Sinorhizobium meliloti (SEQ ID NO: 15); Bradorhizobium japonicum (SEQ ID NO: 16); and Burkholderia fungorum (SEQ ID NO: 17).

Although the degree of sequence identity between the antigens of the invention is low, an appreciation of the antigens at a level beyond simple primary sequence information shows that they share a common arrangement of domains from N-terminus to C-terminus, namely:

- a leader peptide
- a globular head
- a coiled-coil region
- a transmembrane anchor region

Sequence similarity between the various antigens is largely restricted to the C-terminal anchor region. This arrangement of domains is shared with *N.meningitidis* protein NadA {1}.

10 The positions of these features in SEQ ID NO^S: 1 to 18 are as follows:

SEQ ID	Organism	Length	Leader	Head	Coiled-coil	Anchor
1	H.aegyptius	>223	1-26	27-55	56-184	185
2	EHEC	338	1-23	24-207	208-266	267-338
3	Effec	1588	1-53	54-1	515 *	1516-1588
4	A.actinomycetemcomitans	295	1-25	26-150	151-222	223-295
5	H.somnus	452	1-26	27-158	159-378	379-452
6	H.ducreyi	273	1-21	22-1	198 *	199-273
7	EPEC	338	1-24	25-209	210-266	267-338
8	EAEC	717	1-23	24-109	110-645	646-717
9	EAEC	1743	1-53	54-1	670 *	1671-1743
10	UPEC	1778	1-53	1-53 54-1705 *		1706-1778
11	S.flexneri	990		1-917 *		918-990
12	B.melitensis	227	1-27	28-122	123-154	155-227
13	B.suis	311	1-27	28-206	207-238	239-311
14	R.solanacearum	1309	1-2	30 *	231-708	1239-1309
15	S.meliloti	1291	1-1219 *		1220-1291	
16	B.japonicum	372	1-72 73-300 °		300 *	301-372
17	B.fungorum	3399	1-57	58-3	328 *	3329-3399
18	EPEC	577		1-504 *		505-577
51	H.aegyptius	256	1-26	27-55	56-184	185-256

^{*} The boundary between domains is less distinct for some polypeptides of the invention

Antigens

The invention provides a polypeptide comprising one or more of the following amino acid sequences: any of SEQ ID NO^S: 1 to 18, SEQ ID NO: 51, and SEQ ID NO: 54.

The invention also provides a polypeptide comprising an amino acid sequence: (a) having at least m% identity to one or more of SEQ ID NO^S: 1-18, 51 & 54, where m is 50 or more (e.g. 60, 65, 70, 75, 80, 85, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 99.5 or more); and/or (b) which is a fragment of at least n consecutive amino acids of one or more of SEQ ID NO^S: 1-18, 51 & 54, wherein n is 7 or

more (e.g. 8, 10, 12, 14, 16, 18, 20, 25, 30, 35, 40, 50, 60, 70, 80, 90, 100, 150, 200, 250 or more). These polypeptides include variants (e.g. allelic variants, homologs, orthologs, paralogs, mutants, etc.) of SEQ ID NO^S: 1-18, 51 & 54.

Preferred fragments of (b) comprise an epitope from one or more of SEQ ID NO^S: 1-18, 51 & 54, preferably a B-cell epitope. B-cell epitopes can be identified empirically or can be predicted algorithmically.

Other preferred fragments of (b) lack one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25 or more) from the C-terminus and/or one or more amino acids (e.g. 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25, 45 or more) from the N-terminus of the relevant amino acid sequence from SEQ ID NO^S: 1-18, 51 & 54. In particular, preferred fragments omit at least the N-terminus leader sequence (and the omitted leader sequence may be replaced by a heterologous leader sequence).

Other preferred fragments omit one or more (i.e. 1, 2, or 3) of the four domains of SEQ ID NO^S: 1-18 & 51, based on the above table. Other preferred fragments consist of one or more (i.e. 1, 2, or 3) of the four domains of SEQ ID NO^S: 1-18 & 51.

Preferred polypeptides of the invention are presented in oligomeric form (e.g. dimers, trimers, tetramers, etc.). Trimers are preferred, but monomeric polypeptides of the invention are also useful.

The invention also provides polypeptides of the formula $NH_2-A-\{-X-L-\}_x-B-COOH$, wherein:

- X comprises an amino acid sequence: (a) having at least m% identity to one or more of SEQ ID NO^S: 1-18, 51 & 54; and/or (b) which is a fragment of at least n consecutive amino acids of one or more of SEQ ID NO^S: 1-18, 51 & 54, as defined above;
- L is an optional linker amino acid sequence;

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- A is an optional N-terminal amino acid sequence;
- B is an optional C-terminal amino acid sequence; and
- x is 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17 or 18 (preferably x=2).
- Where a -X- moiety has a leader peptide, this may be included or omitted in the hybrid protein. In some embodiments, the leader peptides will be deleted except for that of the -X- moiety located at the N-terminus of the hybrid protein *i.e.* the leader peptide of X_1 will be retained, but the leader peptides of X_2 ... X_x will be omitted. This is equivalent to deleting all leader peptides and using the leader peptide of X_1 as moiety -A-.
- For each x instances of {-X-L-}, -X- may be the same or different, and linker amino acid sequence -L- may be present or absent. For instance, when x=2 the hybrid may be NH₂-X₁-L₁-X₂-L₂-COOH, NH₂-X₁-X₂-COOH, NH₂-X₁-X₂-COOH, NH₂-X₁-X₂-COOH, etc. Linker amino acid sequence(s) -L- will typically be short (e.g. 20 or fewer amino acids i.e. 19, 18, 17, 16, 15, 14, 13, 12, 11, 10, 9, 8, 7, 6, 5, 4, 3, 2, 1). Examples comprise short peptide sequences which facilitate cloning,

poly-glycine linkers (i.e. comprising Gly_n where n = 2, 3, 4, 5, 6, 7, 8, 9, 10 or more), and histidine tags (i.e. His_n where n = 3, 4, 5, 6, 7, 8, 9, 10 or more). Other suitable linker amino acid sequences will be apparent to those skilled in the art. A useful linker is GSGGGG (SEQ ID NO: 19), with the Gly-Ser dipeptide being formed from a *BamHI* restriction site, thus aiding cloning and manipulation, and the $(Gly)_4$ tetrapeptide being a typical poly-glycine linker.

-A- is an optional N-terminal amino acid sequence. This will typically be short (e.g. 40 or fewer amino acids i.e. 39, 38, 37, 36, 35, 34, 33, 32, 31, 30, 29, 28, 27, 26, 25, 24, 23, 22, 21, 20, 19, 18, 17, 16, 15, 14, 13, 12, 11, 10, 9, 8, 7, 6, 5, 4, 3, 2, 1). Examples include leader sequences to direct protein trafficking, or short peptide sequences which facilitate cloning or purification (e.g. histidine tags i.e. Hish where h = 3, 4, 5, 6, 7, 8, 9, 10 or more). Other suitable N-terminal amino acid sequences will be apparent to those skilled in the art. If X_1 lacks its own N-terminus methionine, -A- is preferably an oligopeptide (e.g. with 1, 2, 3, 4, 5, 6, 7 or 8 amino acids) which provides a N-terminus methionine.

-B- is an optional C-terminal amino acid sequence. This will typically be short (e.g. 40 or fewer amino acids i.e. 39, 38, 37, 36, 35, 34, 33, 32, 31, 30, 29, 28, 27, 26, 25, 24, 23, 22, 21, 20, 19, 18, 17, 16, 15, 14, 13, 12, 11, 10, 9, 8, 7, 6, 5, 4, 3, 2, 1). Examples include sequences to direct protein trafficking, short peptide sequences which facilitate cloning or purification (e.g. comprising histidine tags i.e. Hish where h = 3, 4, 5, 6, 7, 8, 9, 10 or more), or sequences which enhance protein stability. Other suitable C-terminal amino acid sequences will be apparent to those skilled in the art.

20 The invention also provides polypeptides comprising the amino acid sequence:

$$-A-W_1-W_2-W_3-W_4-B-$$

wherein:

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- A is an optional sequence as defined above (preferably at the N-terminus of the polypeptide);
- B is an optional sequence as defined above (preferably at the C-terminus of the polypeptide);
- W₁ is an optional amino acid sequence: (a) having at least m% identity to the leader peptide of one or more of SEQ ID NO^S: 1-18 & 51; and/or (b) which is a fragment of at least n consecutive amino acids of the leader peptide of one or more of SEQ ID NO^S: 1-18 & 51;
- W₂ is an optional amino acid sequence: (a) having at least m% identity to the globular head domain of one or more of SEQ ID NO^S: 1-18 & 51; and/or (b) which is a fragment of at least n consecutive amino acids of the globular head domain of one or more of SEQ ID NO^S: 1-18, & 51;
- W₃ is an optional amino acid sequence: (a) having at least m% identity to the coiled-coil domain of one or more of SEQ ID NO^S: 1-18 & 51; and/or (b) which is a fragment of at least n consecutive amino acids of the coiled-coil domain of one or more of SEQ ID NO^S: 1-18 & 51;

- W₄ is an optional amino acid sequence: (a) having at least m% identity to the transmembrane anchor region of one or more of SEQ ID NO^S: 1-18 & 51; and/or (b) which is a fragment of at least n consecutive amino acids of the transmembrane anchor region of one or more of SEQ ID NO^S: 1-18 & 51;

5 provided that at least one of W₁, W₂, W₃ or W₄ is present.

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The invention also provides a polypeptide comprising a polypeptide as described above, wherein the amino acid sequence of the polypeptide contains one or more amino acid mutations. The mutation(s) preferably result in the reduction or removal of an activity of a polypeptide of the invention which is responsible directly or indirectly for virulence or adhesion. For example, the mutation may inhibit an enzymatic activity or may remove a binding site in the protein. Mutation may involve deletion, substitution, and/or insertion, any of which may be involve one or more amino acids. As an alternative, the mutation may involve truncation.

Mutagenesis of virulence factors is a well-established science for many bacteria {e.g. toxin mutagenesis described in refs. 2 to 8}. Mutagenesis may be specifically targeted to nucleic acid encoding a polypeptide of the invention. Alternatively, mutagenesis may be global or random (e.g. by irradiation, chemical mutagenesis, etc.), which will typically be followed by screening bacteria for those in which a mutation has been introduced into a gene encoding a polypeptide of the invention. Such screening may be by hybridisation assays (e.g. Southern or Northern blots etc.), primer-based amplification (e.g. PCR), sequencing, proteomics, aberrant SDS-PAGE gel migration, etc.

Polypeptides of the invention can be prepared by various means (e.g. recombinant expression, purification from cell culture, chemical synthesis, etc.) and in various forms (e.g. native, fusions, non-glycosylated, lipidated, etc.). They are preferably prepared in substantially pure form (i.e. substantially free from other bacterial or host cell proteins).

Whilst expression of the polypeptides of the invention may take place in the native host, the invention preferably utilises a heterologous host. The heterologous host may be prokaryotic (e.g. a bacterium) or eukaryotic. It is preferably *E.coli*, but other suitable hosts include *Bacillus subtilis*, Vibrio cholerae, Salmonella typhi, Salmonella typhimurium, Neisseria lactamica, Neisseria cinerea, Mycobacteria (e.g. M.tuberculosis), yeasts, etc.

Where a polypeptide of the invention is related to SEQ ID NO: 51, it preferably comprises at least 224 (e.g. 224, 225, 226, 227, 228, 229, 230, 235, 240, 245, 250, 255 or more) amino acids.

The invention also provides an adhesin from Haemophilus aegyptius, wherein the adhesin comprises: (a) amino acid sequence SEQ ID NO: 52; (b) an amino acid sequence having at least m% identity to SEQ ID NO: 52; and/or (c) an amino acid sequence which is a fragment of at least n consecutive amino acids of SEQ ID NO: 52.

Antibodies

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The invention also provides antibodies which bind to polypeptides of the invention.

Antibody of the invention preferably has an affinity for a polypeptide of the invention of at least 10^{-7} M e.g. 10^{-8} M, 10^{-9} M, 10^{-10} M or tighter. Preferred antibodies can block the ability of a polypeptide of the invention to bind to a human cell.

Antibodies of the invention may be polyclonal or monoclonal and may be produced by any suitable means (e.g. by recombinant expression, purification from cell culture, chemical synthesis, etc.) and in various forms (e.g. native, fusions, glycosylated, non-glycosylated, etc.). They are preferably prepared in substantially pure form (i.e. substantially free from other antibodies).

The term "antibody" includes whole antibodies, Fv, scFv, Fc, Fab, F(ab')2, etc.

Antibodies of the invention may include a label. The label may be detectable directly, such as a radioactive or fluorescent label. Alternatively, the label may be detectable indirectly, such as an enzyme whose products are detectable (e.g. luciferase, \beta-galactosidase, peroxidase, etc.).

Antibodies of the invention may be attached to a solid support.

Antibodies of the invention may be prepared by administering (e.g. injecting) a polypeptide of the invention to an appropriate animal (e.g. a rabbit, hamster, mouse or other rodent).

To increase compatibility with the human immune system, the antibodies may be chimeric or humanized {e.g. refs. 9 & 10}, or fully human antibodies may be used. Because humanized antibodies are far less immunogenic in humans than the original non-human monoclonal antibodies, they can be used for the treatment of humans with far less risk of anaphylaxis. Thus, these antibodies may be preferred in therapeutic applications that involve *in vivo* administration to a human such as, use as radiation sensitizers for the treatment of neoplastic disease or use in methods to reduce the side effects of cancer therapy.

Humanized antibodies may be achieved by a variety of methods including, for example: (1) grafting non-human complementarity determining regions (CDRs) onto a human framework and constant region ("humanizing"), with the optional transfer of one or more framework residues from the non-human antibody; (2) transplanting entire non-human variable domains, but "cloaking" them with a human-like surface by replacement of surface residues ("veneering"). In the present invention, humanized antibodies will include both "humanized" and "veneered" antibodies. {11, 12, 13, 14, 15, 16, 17}. Humanized or fully-human antibodies can also be produced using transgenic animals that are engineered to contain human immunoglobulin loci.

The phrase "constant region" refers to the portion of the antibody molecule that confers effector functions. In chimeric antibodies, mouse constant regions are substituted by human constant regions.

The constant regions of humanized antibodies are derived from human immunoglobulins. The heavy chain constant region can be selected from any of the 5 isotypes: alpha, delta, epsilon, gamma or mu.

Nucleic acids

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The invention also provides nucleic acid encoding the polypeptides of the invention. Furthermore, the invention provides nucleic acid which can hybridise to this nucleic acid, preferably under "high stringency" conditions (e.g. 65°C in a 0.1xSSC, 0.5% SDS solution).

Nucleic acid according to the invention can be prepared in many ways (e.g. by chemical synthesis, from genomic or cDNA libraries, from the organism itself, etc.) and can take various forms (e.g. single stranded, double stranded, vectors, probes, etc.). They are preferably prepared in substantially pure form (i.e. substantially free from other bacterial or host cell nucleic acids).

The term "nucleic acid" includes DNA and RNA, and also their analogues, such as those containing modified backbones (e.g. phosphorothioates, etc.), and also peptide nucleic acids (PNA), etc. The invention includes nucleic acid comprising sequences complementary to those described above (e.g. for antisense or probing purposes).

15 Immunogenic compositions and medicaments

Based on the structural and functional similarities to NadA, which is a good anti-meningococcal immunogen {1}, including their association with virulence, the polypeptides of the invention should also be useful for immunisation purposes.

The invention provides a composition comprising a polypeptide and/or a nucleic acid and/or an antibody of the invention. Compositions of the invention are preferably immunogenic compositions, and are more preferably vaccine compositions. Vaccines according to the invention may either be prophylactic (i.e. to prevent infection) or therapeutic (i.e. to treat infection), but will typically be prophylactic.

The pH of the composition is preferably between 6 and 8, preferably about 7. The pH may be maintained by the use of a buffer. The composition may be sterile and/or pyrogen-free. The composition may be isotonic with respect to humans.

The invention also provides a composition of the invention for use as a medicament. The medicament is preferably able to raise an immune response in a mammal (i.e. it is an immunogenic composition) and is more preferably a vaccine.

The invention also provides the use of one or more (e.g. 2, 3, 4, 5, 6) of the polypeptides of the invention in the manufacture of a medicament for raising an immune response in a mammal. The medicament is preferably a vaccine.

The invention also provides a method for raising an immune response in a mammal comprising the step of administering an effective amount of a composition of the invention. The immune response is preferably protective and preferably involves antibodies and/or cell-mediated immunity. The method may raise a booster response.

The mammal is preferably a human. Where the vaccine is for prophylactic use, the human is preferably a child (e.g. a toddler or infant) or a teenager; where the vaccine is for therapeutic use, the human is preferably a teenager or an adult. A vaccine intended for children may also be administered to adults e.g. to assess safety, dosage, immunogenicity, etc.

These uses and methods are preferably for the prevention and/or treatment of a disease caused by Haemophilus influenzae biogroup aegyptius, Escherichia coli (particularly EHEC, EAEC, ETEC, EPEC and UPEC strains), Actinobacillus actinomycetemcomitans, Haemophilus somnus, Haemophilus ducreyi, Shigella flexneri, Brucella melitensis, Brucella suis, Ralstonia solanacearum, Sinorhizobium meliloti, Bradorhizobium japonicum and Burkholderia fungorum. Thus the invention is suitable for the prevention and/or treatment of diseases including: conjunctivitis, chancroid, purpuric fever, meningitis, pneumonia, epiglottitis, peri-implantitis, periodontal disease, gingivitis, bovine encephalitis, arthritis, myocarditis, diarrhoea, ovine abortion, orchitis, undulant fever, porcine reproductive wastage, brucellosis, etc.

One way of checking efficacy of therapeutic treatment involves monitoring bacterial infection after administration of the composition of the invention. One way of checking efficacy of prophylactic treatment involves monitoring immune responses against the polypeptides after administration of the composition.

Compositions of the invention will generally be administered directly to a patient. Direct delivery may be accomplished by parenteral injection (e.g. subcutaneously, intraperitoneally, intravenously, intramuscularly, or to the interstitial space of a tissue), or by rectal, oral (e.g. tablet, spray), vaginal, topical, transdermal {e.g. see ref. 18} or transcutaneous {e.g. see refs. 19 & 20}, intranasal {e.g. see ref. 21}, ocular, aural, pulmonary or other mucosal administration.

The invention may be used to elicit systemic and/or mucosal immunity.

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Dosage treatment can be a single dose schedule or a multiple dose schedule. Multiple doses may be used in a primary immunisation schedule and/or in a booster immunisation schedule. In a multiple dose schedule the various doses may be given by the same or different routes e.g. a parenteral prime and mucosal boost, a mucosal prime and parenteral boost, etc.

Bacterial infections affect various areas of the body and so the compositions of the invention may be prepared in various forms. For example, the compositions may be prepared as injectables, either as liquid solutions or suspensions. Solid forms suitable for solution in, or suspension in, liquid vehicles prior to injection can also be prepared (e.g. a lyophilised composition). The composition may be

prepared for topical administration e.g. as an ointment, cream or powder. The composition may be prepared for oral administration e.g. as a tablet or capsule, as a spray, or as a syrup (optionally flavoured). The composition may be prepared for pulmonary administration e.g. as an inhaler, using a fine powder or a spray. The composition may be prepared as a suppository or pessary. The composition may be prepared for nasal, aural or ocular administration e.g. as drops. The composition may be in kit form, designed such that a combined composition is reconstituted just prior to administration to a patient. Such kits may comprise one or more antigens in liquid form and one or more lyophilised antigens.

Immunogenic compositions used as vaccines comprise an immunologically effective amount of antigen(s), as well as any other components, as needed. By 'immunologically effective amount', it is meant that the administration of that amount to an individual, either in a single dose or as part of a series, is effective for treatment or prevention. This amount varies depending upon the health and physical condition of the individual to be treated, age, the taxonomic group of individual to be treated (e.g. non-human primate, primate, etc.), the capacity of the individual's immune system to synthesise antibodies, the degree of protection desired, the formulation of the vaccine, the treating doctor's assessment of the medical situation, and other relevant factors. It is expected that the amount will fall in a relatively broad range that can be determined through routine trials.

The invention also provides the polypeptides of the invention (including NadA itself) for use as adjuvants (parenteral and/or mucosal). Similarly, the invention provides a composition comprising a polypeptide of the invention in admixture with a second antigen, whereby the polypeptide of the invention enhances the immune response against the second antigen when administered to a patient.

Further components of the composition

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The composition of the invention will typically, in addition to the components mentioned above, comprise one or more 'pharmaceutically acceptable carriers', which include any carrier that does not itself induce the production of antibodies harmful to the individual receiving the composition. Suitable carriers are typically large, slowly metabolised macromolecules such as proteins, polysaccharides, polylactic acids, polyglycolic acids, polymeric amino acids, amino acid copolymers, and lipid aggregates (such as oil droplets or liposomes). Such carriers are well known to those of ordinary skill in the art. The vaccines may also contain diluents, such as water, saline, glycerol, etc. Additionally, auxiliary substances, such as wetting or emulsifying agents, pH buffering substances, and the like, may be present. A thorough discussion of pharmaceutically acceptable excipients is available in reference 22.

Vaccines of the invention may be administered in conjunction with other immunoregulatory agents. In particular, compositions will usually include an adjuvant. Preferred further adjuvants include, but are not limited to: (A) aluminium salts, including hydroxides (e.g. oxyhydroxides), phosphates (e.g. hydroxyphoshpates, orthophosphates), sulphates, etc. {e.g. see chapters 8 & 9 of ref. 23}), or mixtures of different aluminium compounds, with the compounds taking any suitable form (e.g. gel,

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crystalline, amorphous, etc.), and with adsorption being preferred; (B) MF59 (5% Squalene, 0.5% Tween 80, and 0.5% Span 85, formulated into submicron particles using a microfluidizer) {see Chapter 10 of 23; see also ref. 24}; (C) liposomes {see Chapters 13 and 14 of ref. 23}; (D) ISCOMs {see Chapter 23 of ref. 23}, which may be devoid of additional detergent {25}; (E) SAF, containing 10% Squalane, 0.4% Tween 80, 5% pluronic-block polymer L121, and thr-MDP, either microfluidized into a submicron emulsion or vortexed to generate a larger particle size emulsion {see Chapter 12 of ref. 23}; (F) RibiTM adjuvant system (RAS), (Ribi Immunochem) containing 2% Squalene, 0.2% Tween 80, and one or more bacterial cell wall components from the group consisting of monophosphorylipid A (MPL), trehalose dimycolate (TDM), and cell wall skeleton (CWS), preferably MPL + CWS (DetoxTM); (G) saponin adjuvants, such as QuilA or QS21 {see Chapter 22 of ref. 23}, also known as StimulonTM {26}; (H) chitosan {e.g. 27}; (I) complete Freund's adjuvant (CFA) and incomplete Freund's adjuvant (IFA); (J) cytokines, such as interleukins (e.g. IL-1, IL-2, IL-4, IL-5, IL-6, IL-7, IL-12, etc.), interferons (e.g. interferon-γ), macrophage colony stimulating factor, tumor necrosis factor, etc. {see Chapters 27 & 28 of ref. 23}; (K) monophosphoryl lipid A (MPL) or 3-O-deacylated MPL (3dMPL) {e.g. chapter 21 of ref. 23}; (L) combinations of 3dMPL with, for example, QS21 and/or oil-in-water emulsions {28}; (M) a polyoxyethylene ether or a polyoxyethylene ester {29}; (N) a polyoxyethylene sorbitan ester surfactant in combination with an octoxynol {30} or a polyoxyethylene alkyl ether or ester surfactant in combination with at least one additional non-ionic surfactant such as an octoxynol {31}; (N) a particle of metal salt {32}; (O) a saponin and an oil-in-water emulsion {33}; (P) a saponin (e.g. QS21) + 3dMPL + IL-12 (optionally + a sterol) {34}; (Q) E.coli heat-labile enterotoxin ("LT"), or detoxified mutants thereof, such as the K63 or R72 mutants {e.g. Chapter 5 of ref. 35}; (R) cholera toxin ("CT"), or detoxified mutants thereof {e.g. Chapter 5 of ref. 35}; (S) double-stranded RNA; (T) microparticles (i.e. a particle of \sim 100nm to \sim 150 μ m in diameter, more preferably \sim 200nm to \sim 30 μ m in diameter, and most preferably ~500nm to ~10µm in diameter) formed from materials that are biodegradable and non-toxic (e.g. a poly(α-hydroxy acid), a polyhydroxybutyric acid, a polyorthoester, a polyanhydride, a polycaprolactone, etc.), with poly(lactide-co-glycolide) being preferred, optionally treated to have a negatively-charged surface (e.g. with SDS) or a positively-charged surface (e.g. with a cationic detergent, such as CTAB); (U) oligonucleotides comprising CpG motifs i.e. containing at least one CG dinucleotide, with 5-methylcytosine optionally being used in place of cytosine; (V) monophosphoryl lipid A mimics, such as aminoalkyl glucosaminide phosphate derivatives e.g. RC-529 {36}; (W) polyphosphazene (PCPP); (X) a bioadhesive {37} such as esterified hyaluronic acid microspheres {38} or a mucoadhesive selected from the group consisting of cross-linked derivatives of poly(acrylic acid), polyvinyl alcohol, polyvinyl pyrollidone, polysaccharides and carboxymethylcellulose; or (Y) other substances that act as immunostimulating agents to enhance the 35 effectiveness of the composition {e.g. see Chapter 7 of ref. 23}. Aluminium salts and MF59 are preferred adjuvants for parenteral immunisation. Mutant toxins are preferred mucosal adjuvants.

Muramyl peptides include N-acetyl-muramyl-L-threonyl-D-isoglutamine (thr-MDP), N-acetyl-normuramyl-L-alanyl-D-isoglutamine (nor-MDP), N-acetylmuramyl-L-alanyl-D-isoglutaminyl-L-alanine-2-(1'-2'-dipalmitoyl-sn-glycero-3-hydroxyphosphoryloxy)-ethylamine MTP-PE), etc.

The composition may include an antibiotic.

5 Further antigens

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As well as containing polypeptides of the invention, the compositions of the invention may also include one or more further antigens. Further antigens for inclusion may be, for example:

- a saccharide antigen from *N.meningitidis* serogroup A, C, W135 and/or Y, such as the oligosaccharide disclosed in ref. 39 from serogroup C {see also ref. 40} or the oligosaccharides of ref. 41.
- antigens from Helicobacter pylori such as CagA {42 to 45}, VacA {46, 47}, NAP {48, 49, 50}, HopX {e.g. 51}, HopY {e.g. 51} and/or urease.
- a saccharide antigen from Streptococcus pneumoniae {e.g. 52, 53, 54}.
- a protein antigen from Streptococcus pneumoniae {e.g. 55}.
- 15 an antigen from hepatitis A virus, such as inactivated virus {e.g. 56, 57}.
 - an antigen from hepatitis B virus, such as the surface and/or core antigens {e.g. 57, 58}.
 - an antigen from hepatitis C virus {e.g. 59}.
 - a diphtheria antigen, such as a diphtheria toxoid {e.g. chapter 3 of ref. 60} e.g. the CRM₁₉₇ mutant {e.g. 61}.
- 20 a tetanus antigen, such as a tetanus toxoid {e.g. chapter 4 of ref. 60}.
 - an antigen from *Bordetella pertussis*, such as pertussis holotoxin (PT) and filamentous haemagglutinin (FHA) from *B.pertussis*, optionally also in combination with pertactin and/or agglutinogens 2 and 3 {e.g. refs. 62 & 63}; whole-cell pertussis antigen may also be used.
 - a saccharide antigen from Haemophilus influenzae B {e.g. 40}.
- 25 polio antigen(s) {e.g. 64, 65} such as OPV or, preferably, IPV.
 - a protein antigen from N. meningitidis serogroup B {e.g. refs. 66 to 77}, such as NadA.
 - an outer-membrane vesicle (OMV) preparation from N.meningitidis serogroup B, such as those disclosed in refs. 78, 79, 80, 81, etc.
 - an antigen from Chlamydia pneumoniae {e.g. refs. 82 to 88}.
- 30 an antigen from Chlamydia trachomatis {e.g. 89}.
 - an antigen from Porphyromonas gingivalis {e.g. 90}.
 - rabies antigen(s) {e.g. 91} such as lyophilised inactivated virus {e.g. 92, RabAvert™}.
 - measles, mumps and/or rubella antigens {e.g. chapters 9, 10 & 11 of ref. 60}.
- influenza antigen(s) {e.g. chapter 19 of ref. 60}, such as the hemagglutinin and/or neuraminidase surface proteins.
 - an antigen from $N.gonorrhoeae \{e.g. 93, 94, 95, 96\}$.

- antigen(s) from a paramyxovirus such as respiratory syncytial virus (RSV {97, 98}) and/or parainfluenza virus (PIV3 {99}).

- an antigen from Moraxella catarrhalis {e.g. 100}, such as UspA1 and/or UspA2
- an antigen from Streptococcus pyogenes (group A streptococcus) {e.g. 101, 102, 103}.
- 5 an antigen from Streptococcus agalactiae (group B streptococcus) {e.g. 104}.
 - an antigen from Staphylococcus aureus {e.g. 105}.
 - an antigen from Bacillus anthracis {e.g. 106, 107, 108}.
 - an antigen from a virus in the flaviviridae family (genus flavivirus), such as from yellow fever virus, Japanese encephalitis virus, four serotypes of Dengue viruses, tick-borne encephalitis virus, West Nile virus.
 - an antigen from Pseudomonas.

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- an antigen from a HIV e.g. a HIV-1 or HIV-2.
- an antigen from a rotavirus.
- a pestivirus antigen, such as from classical porcine fever virus, bovine viral diarrhoea virus, and/or border disease virus.
- a parvovirus antigen e.g. from parvovirus B19.
- a coronavirus antigen e.g. from the SARS coronoavirus.
- a cancer antigen, such as those listed in Table 1 of ref. 109 or in tables 3 & 4 of ref. 110.

The composition may comprise one or more of these further antigens. It is preferred that combinations of antigens should be based on shared characteristics e.g. antigens associated with respiratory diseases, antigens associated with enteric diseases, antigens associated with sexually-transmitted diseases, etc.

Where a saccharide or carbohydrate antigen is used, it is preferably conjugated to a carrier protein in order to enhance immunogenicity {e.g. refs. 111 to 120}. Preferred carrier proteins are bacterial toxins or toxoids, such as diphtheria or tetanus toxoids. The CRM₁₉₇ diphtheria toxoid is particularly preferred {121}. Other carrier polypeptides include the N.meningitidis outer membrane protein {122}, synthetic peptides {123, 124}, heat shock proteins {125, 126}, pertussis proteins {127, 128}, protein D from H.influenzae {129}, cytokines {130}, lymphokines {130}, hormones {130}, growth factors {130}, toxin A or B from C.difficile {131}, iron-uptake proteins {132}, etc. Where a mixture comprises capsular saccharides from both serogroups A and C, it may be preferred that the ratio (w/w) of MenA saccharide:MenC saccharide is greater than I (e.g. 2:1, 3:1, 4:1, 5:1, 10:1 or higher). Different saccharides can be conjugated to the same or different type of carrier protein. Any suitable conjugation reaction can be used, with any suitable linker where necessary.

Toxic protein antigens may be detoxified where necessary e.g. detoxification of pertussis toxin by chemical and/or genetic means {63}.

Where a diphtheria antigen is included in the composition it is preferred also to include tetanus antigen and pertussis antigens. Similarly, where a tetanus antigen is included it is preferred also to include diphtheria and pertussis antigens. Similarly, where a pertussis antigen is included it is preferred also to include diphtheria and tetanus antigens.

Antigens in the composition will typically be present at a concentration of at least lµg/ml each. In general, the concentration of any given antigen will be sufficient to elicit an immune response against that antigen.

As an alternative to using protein antigens in the composition of the invention, nucleic acid encoding the antigen may be used {e.g. refs. 133 to 141}. Protein components of the compositions of the invention may thus be replaced by nucleic acid (preferably DNA e.g. in the form of a plasmid) that encodes the protein.

Processes

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The invention also provides a process for producing a polypeptide of the invention, comprising the step of culturing a host cell transformed with nucleic acid of the invention under conditions which induce polypeptide expression.

The invention provides a process for producing a polypeptide of the invention, comprising the step of synthesising at least part of the polypeptide by chemical means.

The invention provides a process for producing nucleic acid of the invention, comprising the step of amplifying nucleic acid using a primer-based amplification method (e.g. PCR).

The invention provides a process for producing nucleic acid of the invention, comprising the step of synthesising at least part of the nucleic acid by chemical means.

The invention also provides a process for detecting the presence of a bacterium in a sample, comprising the step of contacting the sample with nucleic acid of the invention under hybridizing conditions; and (b) detecting the presence or absence of hybridization of nucleic acid of the invention to nucleic acid present in the sample. The presence of hybridization in step (b) indicates that the sample contains the relevant bacterium.

The invention also provides an immunoassay method for detecting the presence of a bacterium, comprising the step of contacting a sample with a polypeptide or antibody of the invention.

Adhesion inhibition

The invention provides methods for inhibiting the attachment of bacterial cells to host cells (e.g. human cells). The cell may be in vitro (e.g. in cell culture) or in vivo. The cells are most preferably human cells. The host cells will typically be epithelial or endothelial cells.

The invention provides a method for preventing the attachment of a bacterial cell to a host cell, wherein the ability of one or more of the polypeptides of the invention to bind to the host cell is blocked.

The ability to bind may be blocked in various ways but, most conveniently, an antibody specific for a polypeptide of the invention is used. As an alternative to using antibodies, antagonists of the interaction between the polypeptide of the invention and its receptor on the host cell may be used. As a further alternative, a soluble form of the host cell receptor may be used as a decoy. These can be produced by removing the receptor's transmembrane and, optionally, cytoplasmic regions.

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The antibodies, antagonists and soluble receptors of the invention may be used as medicaments to prevent the attachment of a bacterial cell to a host cell.

The invention provides a method for preventing the attachment of a bacterial cell to a host cell, wherein expression of a polypeptide of the invention is inhibited. The inhibition may be at the level of transcription and/or translation. A preferred technique for inhibiting expression of the gene is antisense {e.g. refs. 142 to 148, etc.}. Antibacterial antisense techniques are disclosed in, for example, references 149 & 150.

The invention provides a method for preventing the attachment of a bacterial (e.g. Neisserial) cell to an epithelial cell, wherein the gene encoding the polypeptide of the invention is knocked out. Thus the invention provides a bacterium in which such genes have been knocked out. Techniques for producing knockout bacteria are well known. The knockout mutation may be situated in the coding region of the gene or may lie within its transcriptional control regions (e.g. within its promoter). The knockout mutation will reduce the level of mRNA encoding a polypeptide of the invention to <1% of that produced by the wild-type bacterium e.g. <0.5%, <0.1%, 0%. The knockout mutants of the invention may be used as immunogenic compositions (e.g. as vaccines). Such a vaccine may include the mutant as a live attenuated bacterium.

The invention also provides methods for screening compounds to identify those (antagonists) which inhibit the binding of a bacterial cell to a host cell.

Potential antagonists for screening include small organic molecules, peptides, peptoids, polypeptides, lipids, metals, nucleotides, nucleosides, polyamines, antibodies, and derivatives thereof. Small organic molecules have a molecular weight between 50 and about 2,500 daltons, and most preferably in the range 200-800 daltons. Complex mixtures of substances, such as extracts containing natural products, compound libraries or the products of mixed combinatorial syntheses also contain potential antagonists.

Typically, a polypeptide of the invention is incubated with a host cell and a test compound (e.g. an antibody), and the mixture is then tested to see if the interaction between the protein and the epithelial cell has been inhibited. The protein, cell and compound may be mixed in any order.

Inhibition will, of course, be determined relative to a standard (e.g. the native protein/cell interaction). Preferably, the standard is a control value measured in the absence of the test compound. It will be appreciated that the standard may have been determined before performing the method, or may be determined during or after the method has been performed. It may also be an absolute standard.

For preferred high-throughput screening methods, all the biochemical steps for this assay are performed in a single solution in, for instance, a test tube or microtitre plate, and the test compounds are analysed initially at a single compound concentration. For the purposes of high throughput screening, the experimental conditions are adjusted to achieve a proportion of test compounds identified as "positive" compounds from amongst the total compounds screened.

The method may also simply involve incubating one or more test compound(s) with a polypeptide of the invention and determining if they interact. Compounds that interact with the protein can then be tested for their ability to block an interaction between the protein and an epithelial cell.

Other methods which may be used include, for example, reverse two hybrid screening {151} in which the inhibition of the bacteria:host receptor interaction is reported as a failure to activate transcription.

The invention also provides a compound identified using these methods. These can be used to treat or prevent bacterial infection. The compound preferably has an affinity for a polypeptide of the invention of at least 10⁻⁷ M e.g. 10⁻⁸ M, 10⁻⁹ M, 10⁻¹⁰ M or tighter.

20 **Definitions**

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The term "comprising" encompasses "including" as well as "consisting" e.g. a composition "comprising" X may consist exclusively of X or may include something additional e.g. X + Y.

The term "about" in relation to a numerical value x means, for example, $x\pm 10\%$.

References to a percentage sequence identity between two amino acid sequences means that, when aligned, that percentage of amino acids are the same in comparing the two sequences. This alignment and the percent homology or sequence identity can be determined using software programs known in the art, for example those described in section 7.7.18 of reference 152. A preferred alignment is determined by the Smith-Waterman homology search algorithm using an affine gap search with a gap open penalty of 12 and a gap extension penalty of 2, BLOSUM matrix of 62. The Smith-Waterman homology search algorithm is disclosed in reference 153.

BRIEF DESCRIPTION OF THE DRAWINGS

Figures 1 to 15 show analyses of amino acid sequences of the invention to show coiled-coil regions.

Figure 16 shows conservation between anchor regions of polypeptides of the invention.

Figure 17 is an illustration of the NadA structure within the meningococcal outer membrane, in monomeric and trimeric form.

Figures 18 & 19 show comparisons of the genetic environment of genes encoding polypeptides of the invention. Figure 20 illustrates the genetic environment in *E.coli* K1 vs. K12.

- 5 Figure 21 shows coil analysis for (21A) NadA and (21B) HadA.
 - Figure 22 is a schematic organization of the hadA locus in a hadA positive strain (F3031) and in diverse hadA negative strains (type d, type b, and several non-typeable H. influenzae).
 - Figure 23 is a tree showing the relationship between HadA of different strains.
 - Figure 24 illustrates three constructs for expression of HadA in E.coli.
- Figure 25 shows Bis-Tris gels of expressed HadA-His. Lanes are paired as (odd) total protein and (even) soluble proteins. Lanes 1/2 are empty plasmid at 20°C; 3/4 are expression at 20°C; 5/6 are empty plasmid at 30°C; 7/8 are expression at 30°C; 9/10 are empty plasmid at 37°C; 11/12 are expression at 37°C; M is pre-stained protein standard (See BlueTMPlus2, Invitrogen).
- Figure 26 is a western blot. Lanes are: (1) Pre-stained protein standard, See BlueTMPlus2; (2) empty plasmid, total protein, 30°C; (3) empty plasmid, soluble protein, 30°C; (4) expressed total protein, 30°C; (5) expressed soluble protein, 30°C; (6) rHad A-His.
 - Figure 27 shows FACS analysis of binding to Chang cells by *E.coli*-expressed HadA. HadA was tested at nine concentrations and binding was assessed. Four representative FACS spectra are shown.
- Figure 28 shows phase contrast micrographs of three different aggregates in panels A to C, and cells containing empty pET plasmid in panel D.
 - Figure 29 shows (A) adhesion and (B) invasion of Chang cells by *E.coli* expressing HadA. The left bar is control cells transformed with empty plasmid; the right bar is the HadA-expressing bacteria. Results are the mean \pm standard error of the mean of measurements made in triplicate.
- Figure 30 shows immunofluorescence microscopy analysis of with E. coli-pET HadA na and Chang epithelial cells. Extracellular bacteria are seen in green; intracellular bacteria are red.
 - Figure 31 shows SDS-PAGE of HadA/na and HadA/LNadA/na expressed in *E.coli*. Lanes are: (1) Pre-stained protein standard, See BlueTMPlus2; (2) empty plasmid, overnight uninduced culture; (3) HadA/na, overnight culture; (4) HadA/LNadA/na, overnight culture; (5)-(7) as for (2) to (4), but 3 hours after induction of protein expression by IPTG. Arrows show monomer and oligomer.
- Figure 32 shows a western blot of HadA/na and HadA/LNadA/na expressed in *E.coli*. Lanes are the same as in Figure 31. Arrows show monomer and oligomer.
 - Figure 33 shows SDS-PAGE of HadA expressed overnight in *E.coli* without induction. Lanes are: (1) Pre-stained protein standard, See BlueTMPlus2; (2) empty plasmid; (3) HadA/na-transformed; (4) empty plasmid, outer membrane extract; (5) HadA/na-transformed, outer membrane extract.

Figure 34 shows FACS analysis of HadA expression, showing *E.coli* transformed with an empty pET plasmid or with the HadA/na pET plasmid.

Figure 35 shows the results of a settling assay in E.coli, with or without HadA expression.

MODES FOR CARRYING OUT THE INVENTION

5 Neisseria meningitidis NadA protein

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Within the Neisseria meningitidis serogroup B genome {75}, an outer membrane protein (NadA) was identified {1} which shows weak homology to Yersinia enterocolitica adhesin YadA and to Moraxella catarrhalis surface protein UspA2 {154}. The nadA gene is present in a subgroup of hypervirulent N.meningitidis strains and is characterized by a low GC content, which suggests a probable acquisition event of the gene by horizontal transfer.

To investigate the possibility that proteins similar to the NadA adhesin could have been acquired by other pathogens, we searched for homologous proteins.

A sequence alignment of NadA & YadA revealed that the two proteins are most similar at the C-terminus, which is the membrane anchor domain. In NadA, this domain is approximately 70 residues long and contains five predicted amphipatic beta strands, which cross the outer membrane multiple times thus anchoring the protein to the surface of the bacterium (Figure 17). Within this region, the level of sequence similarity between NadA & YadA is around 60% identity while in the N-terminal and central domain the homology is below 25% identity.

In a first search, based on the NadA anchor domain, results included YadA and UspA2, but also other proteins, such as the serum resistance protein DsrA of *Haemophilus ducreyi*, the immunoglobulin binding proteins EibA-C-D-E and F of *E.coli*, and the outer membrane protein 100 of *Actinobacillus actinomycetemcomitans* {154}. In order to highlight more distant members of this family, these results were used for further searches, and this approach identified 16 further results. These 16 polypeptides were further evaluated for secondary structure analysis, coiled coil prediction and presence/absence of a leader peptide. As expected, despite the little amino acid similarity displayed within the central regions, most of the identified polypeptides possess the coiled coil feature, which gives them the capability to form stable oligomers. The anchor regions of the identified polypeptides are well conserved (Figure 16). In addition, the GC content of the genes encoding these polypeptides was lower than average for their respective genomes, suggesting that they are encoded by genes carried on mobile genetic elements.

Escherichia coli

Polypeptides were found in pathogenic strains of *E.coli*, including enteropathogenic (EPEC), enteroaggregative (EAEC), enterohemorragic (EHEC) and uropathogenic (UPEC) strains. Furthermore, a polypeptide almost identical to those of the EHEC and EPEC strains was found in the

K1 strain, which is a capsulated *E.coli* strain responsible for neonatal meningitis. The K1 sequence aligns with NadA as follows:

```
100
                 110
                          120
                                    130
                                              140
                                                        150
        TGVVQIPARYQSMINARQSAVTDAQQTQITEQQAQIVATQKTLAATGDTQNTAHYQEMIN
k1.pep
                                      1 ::: | :: | | :: | | : | : | : |
NadA.pep DAALADTDAALDETTNALNKLGENITTFAEETKTNIVKIDEKLEAVADTVD--KHAEAFN
         130
                   140
                            150
                                      160
                                                170
                                                           180
       160
                                              200
                 170
                          180
                                    190
                                                        210
k1.pep
         ARLAAQNEANQRTTTEQGQKMNALTTDVAAQQQKERAQYDKQMQSLAQKSVQAHEQIESL
                                :::|::::
                                                1::1:
             :::: | :::: |
NadA.pep DIADSLDETN--TKADEAVKTANEAKQTAEETKQNVDAKVKAAETAAGKAEAAAGTANTA
           190
                      200
                                210
                                          220
                                                    230
                                                             240
       220
                          240
                                    250
                                              260
                                                        270
                 230
         RQDSAQTQQQLTNTQKRVADNSQQINTLNNHFDSLKNEVEDNRKEANAGTASAIAIASQP
kl.pep
          : ::||| ::| : |||:
Nada.pep ADKAEAVAAKVTDIKADIATNKADIAKNSARIDSLDKNVANLRKETRQGLAEQAALSGLF
             250
                      260
                                270
                                          280
                                                    290
                                                             300
       280
                                              320
                 290
                          300
                                    310
                                                        330
         QVKTGDVMMVSAGAGTFNGESAVSVGTSFNAGTHTVLKAGISADTQSDFGAGVGVGYSF
kl.pep
                : 1:1::1 :::1111::11:1
                                         NadA.pep QPYNVGRFNVTAAVGGYKSESAVAIGTGFRFTENFAAKAGVAVGTSSGSSAAYHVGVNYEW
             310
                      320
                                                             360
                                330
                                          340
                                                    350
```

24.4% identity in 209 aa overlap

Another NadA analogue was encoded by the large virulence plasmid present in shiga toxigenic strains of *E.coli* (STEC) {155}. This protein (Saa) is expressed on the outer membrane of *E.coli* and forms high molecular weight oligomers. In contrast, no counterpart of NadA could be detected in the benign *E.coli* strain K12, supporting the view that these genes have been acquired by lateral exchange early during evolution of the species (Figure 20). Nor could a counterpart be seen in laboratory strain MG1655.

Prompted by these observations, and in order to assess a possible mechanism of insertion/deletion of these genes, the arrangement of the region that harbours the gene coding for the NadA-like molecule was investigated. The sequence of this region for the EHEC strain is SEQ ID NO: 23

This analysis showed that the gene organisation of the DNA segments is almost identical among the genomes of K1, EHEC and EPEC, with a sequence conservation of the NadA-like proteins that ranges from 95% identity between K1 and EHEC to 98% identity between K1 and EPEC. In the case of EAEC, although the flanking regions are conserved, the sequence of the NadA-like protein is 380 residues longer than the others, even if the N-terminus and C-terminus are well conserved.

Bacterium	Amino acid	Nucleic acid	Figure
E.coli K1 & E.coli EHEC strain EDL933	SEQ ID NO: 2	SEQ ID NO: 22	3
E.coli EPEC strain E2348/69	SEQ ID NO: 7	SEQ ID NO: 24	
E.coli EAEC strain O42	SEQ ID NO: 8	SEQ ID NO: 25	4

Extending the analysis to the K12 genome, the insertion site was found to be between two hypothetical open reading frames (YbbJ and YbbI) coded on opposite strands, and that the small

"island" consists of three genes: an ORF coding for an hypothetical integral membrane protein, the gene for the putative NadA-like adhesin, and an ORF for a predicted lipoprotein of unknown function. The two latter ORFs are probably co-transcribed, while the first one is coded on the reverse orientation. A couple of 7-bp direct repeats (CTGACGC) that could represent putative insertion sites could be mapped at the boundaries of the inserted fragments (SEQ ID NO: 23, starting at nucleotides 1811 & 4255), and this repeat is absent in the vicinities of the point of insertion in the K12 strain.

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The length of the acquired DNA regions is 2348 bases for EPEC, 2450 bases for K1 and EHEC, and 2630 for EAEC (Figure 18). In all cases, the G+C content of the fragment is lower if compared to the average composition calculated for each genome, thus confirming the preliminary hypothesis that this segment has been acquired by pathogenic *E.coli* by a mechanism of lateral transfer.

In the case of uropathogenic E.coli (UPEC), a different DNA segment was found between the ybbJ ad ybbI genes. This segment is 1342 bp long and encodes a predicted cytoplasmic protein, which is conserved only in Salmonella typhymurium LT2, but absent from all the other analyzed strains of E.coli. Differently from the other described insertion fragments, no direct repeats could be mapped at the boundaries of this island, whose GC composition is also very similar to the average value. These data could indicate that the NadA-like encoding gene has been inserted later on in place of the c0608 gene. Nevertheless, subsequent search revealed that a gene coding for an homologue of NadA could be found in a different location of the genome of uropathogenic E.coli strain CFT073. This protein is more distantly related to NadA and is seen as a member of a second NadA-like family of proteins. Counterparts of this protein are contained in the other pathogenic strains of E.coli at analogous locations and, similarly to the first group of E.coli NadA-like molecules, the corresponding genes are also encoded on small islands and are not present in the K12 strain (Figure 19). Furthermore, these genes have strong similarities at the 3' end with a frame-shifted Shigella flexneri sequence. The arrangement of NLM flanking regions has been compared in the two species (E.coli and Shigella) revealing striking similarities. Although the sequence conservation is restricted to the amino and carboxy-terminal portions of the adhesin coding genes, the flanking regions are syntenic and share more than 80% identity at the nucleotide level. Upstream of the NadA-like gene, this island contains an ORF coding for a lipoprotein that is frameshifted either in EPEC, EHEC and in Shigella. Furthermore, in the genome of Shigella, two additional genes (insA and insB), coding for transposase elements are found in the vicinities of the NLM gene.

Bacterium	Amino acid	Nucleic acid	Figure	
E.coli UPEC strain CFT073	SEQ ID NO: 10	SEQ ID NO: 26	5	
E.coli EHEC	SEQ ID NO: 3	SEQ ID NO: 27	6	
E.coli EAEC	SEQ ID NO: 9	SEQ ID NO: 28	7	
E.coli EPEC	SEQ ID NO: 18	SEQ ID NO: 30	8	
S.flexneri	SEQ ID NO: 11	SEQ ID NO: 31	9	

Haemophilus

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An incomplete NadA homolog was found in Brazilian purpuric fever (BPF) Haemophilus influenzae isolates {156}. This polypeptide has been named HadA. NadA and HadA align as follows:

			10	20	30	40
HadA.pep		MKRNLLK	QSVIAVLIG	GTTVSNYALA	AQAQAQAQVKI	KDELSELKKQVKEM-
				:: 111	:::1 :	: :: ::: :
NadA.pep	-	NVDAKVKAA	ESEIEKLTT		ADTDAALDETI	NALNKLGENITTFA
	100	110	120	130	140	150
		•-				
	50	60	70	80	90	100
HadA.pep			=			AKGDSSEALDKIEAL
77 - A B						
NadA.pep						VKTANEAKQTAEET
	160	170	180	190	200	210
1	10 :	120	130	140	150	160
HadA.pep		• •	F = = =			SADIATNTAAIATH
				1 :: :1		: 1111111 : 111:
NadA.pep				·		CKADIATNKADIAKN
	220	23	0	240	250	260 270
	170	180	190	200	210	220
HadA.pep			-			/GGYKSQTAVAVG
	• • •			• • • • • • • •		
NadA.pep						/GGYKSESAVAIGTG
	280) 2	90	300	310	320 330
NadA.pep	בים בייים בית בית בית	AKAGVAVGT	G G G G G N N V LI	UCINIVEW		
naum.pep	340		50 50	360		
	241	ر ر	J U	300		

No HadA counterpart could be detected either in non-typeable *H.influenzae* strain 86028, which is responsible for otitis media in children, or in the non-pathogenic *H.influenzae* strain Rd KW20. The very high level of sequence identity between HadA and NadA in the C-terminal anchor region might indicate a common origin.

In order to analyze the origin of the hadA gene, the nucleotide sequence of this DNA region in the BPF isolate (SEQ ID NO: 20) was compared to the same region in the genome sequence for *H.influenzae* strains: the non-pathogenic strain Rd {157}, and a non-typeable 86028 strain (NTHi 86028), associated with pediatric otitis media disease.

The results of this comparison indicate that the adhesin coding gene is specific for the Brazilian Purpuric Fever clone (strain F3031), while no counterparts could be mapped either in the laboratory Rd or in the non-typeable strains. The HadA-encoding fragment has an organization that closely resembles that described for NadA {1} and includes an intact open reading frame plus a 182 bp upstream region, which contains -10 and -35 promoter elements. The small genetic island is flanked by the RNA helicase gene at the 5' end and by a putative protease encoding gene located at the 3' end. The GC composition of the recombined segment is consistent with the rest of the genome.

In contrast, while the NTHi 86028 strain can be regarded as a totally negative strain as it lacks the whole region encompassing the RNA helicase and protease ORFs, the Rd genome contains at this location a DNA segment of 1.1 kb, which encodes two short ORFs of unknown function. This region

is characterized by an abnormal GC content (32%) thus suggesting that an independent recombination event has taken place at this site.

Additional NadA-like molecules were identified in other Haemophilus species, namely H.somnus, H.ducreyi and H.actinomycetemcomitans (also known as Actinobacillus actinomycetemcomitans).

Bacterium	Amino acid	Nucleic acid	Figure
H.influenzae biogroup aegyptius	SEQ ID NO: 1	SEQ ID NO: 20	1
H.somnus strain 129PT	SEQ ID NO: 5	SEQ ID NO: 21	2
H.ducreyi	SEQ ID NO: 6	_	_
H.actinomycetemcomitans	SEQ ID NO: 4	_	

NadA and the *H.actinomycetemcomitans* sequence align as follows:

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		^				
	10	20	30	40	5	
actac.pe	MTYQLFKHHLVAL	MVTGAISVNAL	AKDSFLENPS	ANLPQQVFKN : : :	RVDIF :	NNETNI ::
NadA.pep	TIYDIGEDGTITQK	DATAADVEADD		TNLTKTVNEN	KQNVDAKVK	
	60 70	80	90	100	110	
	60 7	0 80	90	100		
actac.pe	NENKKDIAINKANI	ASIEKDVMRNT	GGIDRLAKQE	LVNRARITKN	IELDIRKN T K	SIAENT
31-43	:: : : : EKLTTKLADTDAAL		::::: ::: MATNKT.CEN-	 	; ; ; PARETKTNIV	KIDEKL
NadA.pep	120 130		150	111	160	170
				150	4	CO
	120 1	.30 14		150 - אאג פפאאד-		.60 Tatenka
actac.pe	ASIA-RIDGNLEGV	NKATÕNADAKS	1:			1 1: 1
NadA.pep	EAVADTVDKHAEA	NDIADSLDETN	TKADEAVKT <i>I</i>	NEAKOTAEET	KONVDAKVK	AAETAA
	180	190	200	210	220	230
	170	180	190	200	210	220
actac.pe	DKADVEKNRADIA	N-SRAIAT-FF	RSSSONIAALI	TKVDRNTAR	DRLDSRVNE	LDKEVK
	11:: : 1: []	:::::::	::: :	: : :		
NadA.pep	GKAEAAAGTANTA		OTKADIATNI 260	ADIAKNSAR. 270	280	290
	240	250	200	270	200	
	230	240	250	260	270	280
actac.pe	NGLASQAALSGLF	OPYNVGSLNLS#	AAVGGYKSKT	ALAVGSGYRFI	NONVAAKAG\	/AVSTN-
	:111 11111111			: : : : 	:: 	
NadA.pep	QGLAEQAALSGLF(2PYNVGRFNVTA 310	AVGG1K5E52 320	330	340	350
	500	710				
	290					
actac.pe	GGSATYNVGLNFE					
Nada nen	: : : : GSSAAYHVGVNYE					
naun. pep	360	•		•		
	-			27 NS	identity	in 284 aa 6

37.0% identity in 284 aa overlap

NadA and the *H.somnus* sequence align as follows:

```
160
                                          170
                                                     180
                                                               190
                     150
  H.somnus.pep GF----VEDVNES---EARNISRIDENEKDIKNL--KKELYDFVEDVNESEARNISRID
                    : |::|: :: |: |:|:| || |:::: |||:::
      NadA.pep GFKAGETIYDIGEDGTITQKDATAADVEADDFKGLGLKKVVTNLTKTVNENKQNVDAKVK
                                                90
                                                                 110
                    60
                             70
                                       80
                                                        100
                 200
                           210
                                      220
                                               230
                                                        240
                                                                 250
  H.somnus.pep ENEKDINTLK-ELMDED--LNSVLTQIEDVKLTFQDVNDNVNLAFEEINGNAQKFDTAIE
                ]::[: ] :| ] | ] :: : :::: ::: ::: :::
      NadA.pep AAESEIEKLTTKLADTDAALADTDAALDETTNALNKLGENITTFAEETKTNIVKIDEKLE
                   120
                            130
                                      140
                                               150
                                                        160
                                                                 170
                    260
                             270
                                      280
                                               290
                                                        300
                                                                 310
  H.somnus.pep GLTSGLSDLQAKVDANKQETEDDIADNAKAIHSNTKGIAKNTKDIRDLDTKTKQMLENDK
                         11: 1::1111:
                                          ::::
                                               :|:::: ::
      NadA.pep AVAD-----TVDKHA-EAFNDIADSLDETNTKADEAVKTANEAKQTAEETKQ-----
                                    190
                                             200
                          180
                                                      210
                    320
                             330
                                      340
                                               350
                                                        360
                                                                 370
  H.somnus.pep NLMTGLESLATETSKGFERFDVKTQQLDQAVANVVGRVDITEQAIRQNTAGLVNVNKRVD
              NadA.pep NVDAKVKAAETAAGKAEAAAGTANTAADKAEA-VAAKVTDIKADIATNKADIAKNSARID
                                                   260
              220
                       230
                                240
                                          250
                                                             270
                                            400
                          380
                                   390
                                                      410
                                                               420
  H.somnus.pep TLDKN-----TKAGIASAVALGMLPQSTAPGKSLVSLGVGHHRGQSATAIGVSSMSSN
                     NadA.pep SLDKNVANLRKETRQGLAEQAALSGLFQPYNVGRFNVTAAVGGYKSESAVAIG-TGFRFT
               280
                        290
                                 300
                                          310
                                                    320
                                                              330
                 430
                          440
                                   450
  H.somnus.pep GKWVVKGGMSYDTQRHATFGGSVGFFFN
               NadA.pep ENFAAKAGVAVGTSSGSSAAYHVGVNYEW
                340
                         350
                                  360
                                                  23.2% identity in 354 aa overlap
NadA and the H.ducreyi sequence align as follows:
                  150
                                    170
                                             180
                           160
                                                      190
  H.ducreyi.pe SKNKQNIDTISKYLLELGTYLDGSYRMMEQNTHNINKNTHNINKNTHNINKLSKELQTGL
                                   NadA.pep EAAAGTANTAADKAEAVAAKVTDIKADIATNKADIAKNSARIDSLDKNVANLRKETROGL
                  240
                           250
                                    260
                                             270
                                                                290
                  210
                           220
                                    230
                                             240
                                                      250
                                                                260
  H.ducreyi.pe ANQSALSMLVQPNGVGKTSVSAAVGGYRDKTALAIGVGSRITDRFTAKAGVAFNTYNGG-
              1:1:11 | [] :[]: :[:[]]]]]:::::|:[[]:]: |:[:]:[]:]:
      NadA.pep AEQAALSGLFQPYNVGRFNVTAAVGGYKSESAVAIGTGFRFTENFAAKAGVAVGTSSGSS
                  300
                           310
                                    320
                                             330
                                                      340
                                                                350
                   270
  H.ducreyi.pe MSYGASVGYEF
```

47.5% identity in 101 aa overlap

NB: the coiled-coil prediction for the *H.ducreyi* polypeptide is not high.

Other bacteria

Further NadA homologs identified in the search are:

:|::|::|:

NadA.pep AAYHVGVNYEW

Bacterium	Amino acid	Nucleic acid	Figure	
Brucella melitensis	SEQ ID NO: 12	SEQ ID NO: 32	10	

Brucella suis	SEQ ID NO: 13	SEQ ID NO: 33	11
Ralstonia solanacearum	SEQ ID NO: 14	SEQ ID NO: 34	12
Sinorhizobium meliloti	SEQ ID NO: 15	SEQ ID NO: 35	13
Bradorhizobium japonicum	SEQ ID NO: 16	SEQ ID NO: 36	14
Burkholderia fungorum	SEQ ID NO: 17	SEQ ID NO: 29	15

Multiple sequence alignment

A multiple sequence alignment of members of the NadA "family" is below:

	10	20	30	40	50	60
. 0.51			1		1	1
961_HI	MKRNLLKQ					
961_ACTAC	MTYQLFKH					
MenB NadA	-MSMKHFPSKVLTT					
YADA_YEREN	MTKDFKISVSAA					
961_HAESO 961_K1	MKKVQFFKYSSLAL					
961 HAEDU	MKTVNVALLAL					
301_UMEDO	MKIKCLVA	· A A GTHC S.T.T.	LIMMÖÖB	,		·
Prim.cons.	M23MK42K22LLA2	AI2A2FS2G	ALAA2T6D44	4TGPEA33V3	13P3A333L33	333333
	70	80	90	100	110	120
	1	1	i			1
961_HI		YALAQAQAQA	AQVKKD		, 	
961_ACTAC	AKDSF			VDIFNNET		
MenB NadA	ETIYDIGEDGTITQ	KDATAADVE	ADDFKGLGLK	KVVTNLTK		
YADA_YEREN	PVPGAGGLNASAKG	IHSIAIGAT	AEAAKGAAVA	VGAGSIATGV1	1SV	AIG
961_HAESO	LENEVAYLRMKAGE	WMQLGLDPE	KEVIKGWNEV	KSLPRIDGNG	KDKQTKDQIAM	LIRTVD
961_K1	ELSAIN	SGMSQSEIE	QKITRFLERT	DNSPAAYT		
961_HAEDU	PKFA	GVSSLYSYE	YDYGKGK			~~~~
_			. :		٠	
Prim.cons.	333333GL4A2A66	77SS2ADAE	A3VFKGL444	255PNI5T222	222QTKDQIAM	ILIR222
	130	140	150	160	170	180
	1	1	1	1	1	1
961_HI	ELSELKKQVKEMDA	AIDGILDDN	IAYEAEVDAK	LDQHSAALGRI	TNRLNNLKT-	
961_ACTAC	NINENKKDIAINKA					
MenB NadA	TVNENKQNVDAKVK	AAESEIEKL:	PTKLADTDAA	LADTDAALDET	TNALNKLGE-	
YADA_YEREN	PLSKALGDSAVTYG	AASTAQKDG	VAIGARASTS:	DTGVAVGFNSI	KADAKNSVAIG	HSSHVA
961_HAESO	NTKELGRIVSTNIE	DIKNLKKEL	YGFVEDVNES	EARNISRIDE 1	VEKDIKNLKK~	
961 <u>K</u> 1	YLTEHHYIPSETPD	TTQTPPVQTI	DPDAGQKTVA:	ATGVVQIPAR Y	QSMINARQS~	
961_HAEDU	WTWSNEGGFDIKVP	GIKMKPKEW:	ISKQATYLEL	QHYMPYTPVL1	/TSAPDVSPS-	
Prim.cons.		A TWO TRYET		000::202020		
FIIM. Cons.	NL2ENK22V323VA		LAK /AUVUZ3	222V / ZAZZR :	T3AZNNLKSG	HSSHVA
	190	200	210	220	230	240
			i	•	1	1
961_HI				-IAEKAKGDSS	SEALDKIEALE	EQNDE-
961_ACTAC				-Kntksiaent	'ASIARIDGNL	EGVNR-
MenB NadA	M-H-4					
YADA_YEREN	ANHGYSIAIGDRSK	TDRENSVSI	GHESLNRQLT:	HLAAGTKDTDA	VNVAQLKKEI	EKTQEN
961_HAESO	************		ELY	DFVEDVNESE	ARNISRIDENE	KDINTL
961_K1				AVTDAQQTQII	PEQQAQIVATQ	KTLAAT
961_HAEDU	~			SISILLYPN	ISDPDQLGINR	QQLKLN
Prim.cons.	ANHGYSIAIGDRSK	TDRENSVSI	GHESLNR2L2	36A2K7KEE72	:: ENIAQID2N2	: EQ22E2
	250	260	270	280	290	300
061 117	1	1	1	ı)	70707
	77					
961_ACTAC MenB NadA	UDZUNENEUDTADO					
	VDKHAEAFNDIADS					-
YADA_YEREN	TNKRSAEL					
961_HAESO	KELMDEDLNSVLTQ			· -		
961_Kl	GDTQNTAHYQE					
961_HAEDU	LYSY				LKVLDARI	SKNKQN
Prim.cons.	4DK44E22N34257	: : LA22227A22	25A52VNL22	222222223	TT7N3L2QKI	AE2K2N

	310	320	330	340	350	360
961 HI	ISDIED	<u> </u>		DTN	ן יימייע דמע פארט	ן שיימדממי
961 ACTAC	IENKADKA					
MenB NadA	VDAKVKAA			ETA	agkaeaaagt	ANTAAD
YADA_YEREN	LNMAKAHSNSVART					
961_HAESO 961_K1	VDANKQETEDDIAD VAAQQOKE					
961_HAEDU	IDTISK					
_	:					
Prim.cons.	2DA2K3KA222222	222222222	22A2NTKDI	22L2T223D72	2NSA23AA31	'22IATE
	370	380	390	400	410	420
061 UT	İ	1	1		NOUNDI NEDI	PDCI NA
961_HI 961_ACTAC	RSSSQ			TQRLDNLD ntaridrld		
MenB NadA	KAEAVAAKVTD					
YADA_YEREN	KSSHTLKTANS		_			
961_HAESO	TSKGFERFDVKTQQ					
961_K1 961_HAEDU	QIESLRQDSAQ					
_				:		* *
Prim.cons.	2S22FE4544K44Q	44Q5IANN61	2VA13EQ31	24NTARID2LI)NRVN2LDKE	KAGLAS
	430	440	450	460	470	480
961 HI	QAALNGLFQPYNVG	¦ KT.NT.TAAVGO	YKSOTAVAV(\ 	l
961 ACTAC	QAALSGLFQPYNVG					
MenB NadA	QAALSGLFQPYNVG					
YADA_YEREN	SAALNSLFQPYGVG					
961_HAESO 961 K1	AVALGMLPQSTAPG AIAIASQPQVKTGD					
961_HAEDU	QSALSMLVQPNGVG					
_ •				* :		
Prim.cons.	QAALSGLFQPYNVG	KLNVSAAVG(SY2S32AZAI	G3GS2RFNEN2	ZAAKAGVAZDI	rQZGGSS
	490 !					
961 HI		SEQ ID NO): 1			
961_ACTAC	ATYNVGLNFEW	SEQ ID NO				
MenB NadA	AAYHVGVNYEW	SEQ ID NO			•	
YADA_YEREN 961 HAESO	VMYNASFNIEW FGGSVGFFFN	SEQ ID NO				
961_K1	AGVGVGYSF	SEQ ID NO				
961 HAEDU	7104010101	SEG ID W): Z			
	YGASVGYEF	SEQ ID NO				

HadA studies

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As mentioned above, the HadA sequence (SEQ ID NO: 1) was initially found in an incomplete form. The complete HadA locus was amplified using a forward oligonucleotide primer $HOM\ F$ (SEQ ID NO: 40), a reverse primer $HOM\ R2$ (SEQ ID NO: 41) and an alternative reverse primer $HOM\ R3$ (SEQ ID NO: 42) that is further downstream than $HOM\ R2$. Seven primers (forward: SEQ ID NOS: 43 to 46; reverse: SEQ ID NOS: 47 to 49) were used to sequence the complete locus.

The complete locus in F3031 strain is given as SEQ ID NO: 50. Nucleotides 874 to 1339 of this sequence are new and downstream of SEQ ID NO: 20. The amino acid sequence of HadA is SEQ ID NO: 51. The C-terminus downstream of SEQ ID NO: 20 is given separately as SEQ ID NO: 52.

10 An alignment of NadA and HadA (39.5% identity in 243 aa overlap) is given below:

					10	20	30	40	50	60
HadA			MKRI	NLLKQSVIAVI				KKQVKEM-DAA		
						• • •				
NadA	FKGLGLKK	/VTNLTKTVI	ienkonvdak ⁱ	vkaaeseieki	LTTKLADTDA	ALADTDAAL	DETTNALNKI	Genittfaeet	KTNIVKIDE	KLEAVADT
	80	90	100	110	120	130	140	150	160	170

```
160
                                                                             150
                                                        130
                                                                 140
            80
                             100
                                      110
                                               120
   70
                     90
KLDQHSAALGRHTNRLNNLKTIAEKAKGDSSEALDKIEALEEQNDEFLADITALEEGVDGLDDDITGIQDNISD----IEDDINQNSADIATNTAAIATH
                                           1:11::1::1::1
 :|:|: |:: |:: |:: |::|
                          :: | | : : : |
-VDKHAEAFNDIADSLDETNTKADEAVKTANEAKQTAEETKQNVD---AKVKAAETAA-GKAEAAAGTANTAADKAEAVAAKVTDIKADIATNKADIAKN
                                                                                       270
                                                                     250
                                                                              260
                                                  230
                                                            240
                             210
                                      220
  180
           190
                     200
                                                                               250
                                                                     240
                                                   220
                                                            230
                                 200
                                          210
      170
               180
                        190
TQRLDNLDNRVNNLNKDLKRGLAAQAALNGLFQPYNVGKLNLTAAVGGYKSQTAVAVGTGYRYNENIAAKAGVAF--THGGSATYNVGVNFEW
: 1:11:1:1111:11
SARIDSLDKNVANLRKETRQGLAEQAALSGLFQPYNVGRFNVTAAVGGYKSESAVAIGTGFRFTENFAAKAGVAVGTSSGSSAAYHVGVNYEW
                                                                                360
                                                                        350
                                                               340
                                                      330
                                             320
         280
                  290
                           300
                                    310
```

Although the overall identity is 39.5%, the identity in the C-terminus portion is much higher (up to 86%). Although the central domains of the two proteins are not well conserved, both proteins are predicted to adopt a strong coiled-coil conformation (Figures 21A & 21B).

A schematic organization of the hadA locus in a hadA positive strain (F3031) and in diverse hadA negative strains (type d, type b, and several non-typeable H. influenzae) is shown in Figure 22. The flanking genes are always conserved: they are HI0422, a RNA helicase and HI0419, a putative protease, both in a reverse orientation with respect to hadA.

Immediately downstream of hadA is a gene encoding a hypothetical protein (SEQ ID NOS: 53 & 54), which is frame-shifted in strain KW20 and absent from all other *Haemophilus* strains tested. The closest database match for this protein is ZP_00132218.1, the histone acetyltransferase HPA2 and related acetyltransferases from *Haemophilus somnus* 2336 (SEQ ID NO: 55):

```
Length = 168

Score = 276 bits (707), Expect = 9e-74
Identities = 139/168 (82%), Positives = 149/168 (88%)

Query: 1 MINENLAYLSVLPLEDVKIERSSFSCSVEPLENYFHKYVSQDVKKGLAKCFVLINAQPSR 60
MINENL YLSVLPLEDH I+R+SFSCSVEPLE YF+KY SQDVKKG+ KCFVLIN Q

Sbjct: 1 MINENLPYLSVLPLEDLTIDRNSFSCSVEPLETYFYKYASQDVKKGITKCFVLINKQQFG 60

Query: 61 IVGYYTLSALSIPIPDIPQERISKGVPYPNIPAVLIGRLAIDTNFQKQGYGKFLIADAIH 120
I+GYYTLSALSIPI DIPQERISKG+PYPNIPAVL+GRLAIDTNFQ QGYGKFLIADAI+
Sbjct: 61 IIGYYTLSALSIPITDIPQERISKGIPYPNIPAVLVGRLAIDTNFQNQGYGKFLIADAIY 120

Query: 121 KIKNATVAATILVVEAKNDDASSFYERLGFIEFKEFGGTHRKLFYPLT 168
KIKNATV A ILVVEAKND A SFY+RLGFIEFK THRKLFYPLT
Sbjct: 121 KIKNATVGAAILVVEAKNDHAVSFYKRLGFIEFKNLKKTHRKLFYPLT 168
```

An alignment of the locus in HadA-negative strains is given below:

CLUSTAL W (1.83) multiple sequence alignment

86028	GCAAGCCAAGTAACAGTAATGTTTAATTAGGTATGATTTAAATTCTGTTTTATATCACAC
R2846	GCAAGCCAAGTAACAGTAATGTTTAATTAGGTATGATTTAAATTCTGTTTTATATCACAC
NT36	GCAAGCCAAGTAACAGTAATGTTTAATTAGGTATGATTTAAATTCTGTTTTATATCACAC
EAGAN	GCAAGCCAAGTAACAGTAATGTTTAATTAGGTATGATTTAAATTCTGTTTTATATCACAC
НК707	GCAAGCCAAGTAACAGTAATGTTTAATTAGGTATGATTTAAATTCTGTTTTATATCACAC

WO 2004/113371	PCT/IB2004/002351		
R2866	GCAAGCCAAGTAACAGTAATGTTTAATTAGGTATGATTTAAATTCTGTTTTATATCACAC **************************		
86028	TAGCAATGTGGGTTTCTTGTATTGGTATTAACTAAATTACGCATTAATAAAGCGTAATTT		
R2846	<u>TAGCAATGTGGGTTTCTTGTATTGGTATTAACTAAATTACGCATTAATAAAGCGTAATTT</u>		
NT36	<u>TA</u> GCAATGTGGGTTTCTTGTATTGGTATTAACTAAATTACGCATTAATAAAGCGTAATTT		
EAGAN	TAGCAATGCGGGTTTCTTGTATTGGTATTAACTAAATTACGCATTAATAAAGCGTAATTT		
HK707	TAGCAATGCGGGTTTCTTGTATTGGTATTAACTAAATTACGCATTAATAAAGCGTAATTT		
R2866	TAGAAATGAGGATTTCTTGTATTGGTATTAACTAAATTACGCATTAATAAGGCGTAATTT *** *** ** **********************		
86028	AAGTTAATATCTTGTGGTACATTTAAGAATACAAAATGCCCATCACCTAGTG SEQ ID NO: 56		
R2846	AAGTTAATATCTTGTGGTACATTTAAGAATACAAAATGCCCATCACCTAGTG SEQ ID NO: 57		
NT36	AAGTTAATATCTTGTGGTACATTTAAGAATACAAAATGCCCATCGCCTAGTG SEQ ID NO: 58		
EAGAN	AAGTTAATATCTTGTGGTACATTTAAGAATACAAAATGCCCATCGCCTAGTG SEQ ID NO: 59		
нк707	AAGTTAATATCTTGTGGTACATTTAAGAATACAAAATGCCCATCGCCTAGTG SEQ ID NO: 60		
R2866	AAGTTAATATCTTGTGGCACATTTAAGAATACAAAATGCCCATCGCCTAGTG SEQ ID NO: 61		

The EAGAN and HK707 sequences are from type b Hi strains; the other four are from NTHi strains. The TAA stop codon of the upstream gene (HI0422) is underlined, as is the reverse complement of the TAG stop codon of the downstream gene (HI0419). The HadA gene is seen between these two sequences, and the key intergenic sequence is SEQ ID NO: 62.

Although *H.influenzae* strains Rd and F1947 lack the HadA gene, the sequence between HI0419 and HI0422 is longer, and includes a sequence that has homology to the region upstream of and including the first five codons of HadA. The Hi biogroup aegyptius sequences are as follows:

CCGACGCAAGCCAAGTAATAGTAATATT*TAA*TTAGGTATGATGTAAATTCTGCTTGAGGC end of HI0422 * similar to SEQ ID NO: 62

ACTCAGTTATGGTTATGGTTACATAGTATAGTTTTACTTTGTTCTAGTTCACTTTAATAA

CCTTAAATAATTGAGGATTTCTTATGAAAAGAAATTTATTAAAACAATCTGTAATCGCTG .

M K R N L L K Q S V I A V HadA

TGTTGATAGGTGGCACTACTGTTTCTAATTATGCTTTAGCACAAGCACAAGCACAAGCAC... SEQ ID NO: 64 L I G G T T V S N Y A L A Q A Q A Q ... SEQ ID NO: 1

The underlined 77mer (SEQ ID NO:63) is also seen in strains Rd and F1947, downstream of HI0422:

AGGATACGAAAAATATCGGCAAACGACGCAAGCCAAGTAACAGTAATGTT*TAG*GCTTGTA

TAGTATAGCTTTGCTTCTAGTTCAATTTAATAATCTTAAATAATTAAGGATTTCTT

ATGAAAAAAATTTATAGGCTTCGTTTCGCACACTCGTTGCTAGTATAGATATGTGAATA... SEQ ID NO: 65

This shared sequence could cause some level of cross-reaction in southern blots even though strains may be HadA-negative.

Southern blot experiments on a panel of various *Haemophilus* strains revealed the presence of HadA in a variety of strains (Figure 23). All other typeable and non-typeable strains of *H.influenzae* lacked the *hadA* gene by this analysis.

HadA expression in E.coli

To study the structure and function of HadA, different constructs were prepared for expression in *E.coli*, as illustrated in Figure 24: (1) to express the protein as full length (native HadA, or 'HadA/na'); (2) HadA under the control of NadA leader peptide ('HadA/LNadA/na') and (3) as a C-terminal histidine fusion ('HadA-his'). All the constructs were made in pET21b expression vector and *E.coli* BL21(DE3) was used as expression host.

HadA-his was expressed at different temperatures. Total and soluble proteins were analysed by SDS-PAGE (Bis-Tris gels, 12% MOPS; InvitrogenTM). The gels are shown in Figure 25. The soluble protein expressed at 30°C (lane 9) was purified and was used to immunise mice.

Purification of HadA-His can be performed by the IMAC process, but this was not very efficient for removing all *E.coli* contaminants. IMAC was thus followed by dialysis in a pH 7.7 buffer, for anionic exchange chromatography. Surprisingly the protein was completely precipitated. Subsequently the protein was dialysed in four different pH condition (6.3, 6.5, 7.7 & 8.5) and precipitation was seen only at pH 7.7. With a theoretical pI of 4.38 then the precipitation should not be isoelectric precipitation.

Sera from the mice were used to visualise western blots (12% Mops) of different fractions of *E.coli* strains and purified recombinant HadA. The first antibody was the anti-HadA (1:1000); the second antibody was anti-mouse immunoglobulin-HRP (DAKO) 1:10000. The results are in Figure 26.

SDS-PAGE of HadA/na and HadA/LNadA/na is shown in Figure 31. In both overnight and induced cultures, HadA protein was expressed as monomer and as an oligomer (e.g. trimer) that is heat stable in SDS gel. Western blotting is shown in Figure 32, to confirm the presence of HadA monomer and oligomer in the *E.coli* bacteria.

Expression was also studied by examining bacterial outer membrane proteins. Figure 33 shows SDS-PAGE (Bis-Tris gel 10% MOPS, Invitrogen) of an OMV preparation from *E.coli* and shows that HadA oligomers are seen in the outer membranes. The cell-surface location was confirmed by FACS, as shown in Figure 34.

30 Adhesion

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Purified HadA was tested for its ability to bind to Chang epithelial cells. The experiments showed by FACS analysis that HadA-his binds to the cells in a dose dependent manner (Figure 27).

E.coli BL21(DE3) that express HadA/na were tested for aggregation. Figure 28 shows phase contrast micrographs of cellular aggregates collected from late exponential phase cultures that had been left

standing at room temperature for 4 hours. Three different samples are shown in 28A to 28C. The bacteria form visible bacterial "clouds", and bacterial aggregation can be correlated with microcolony formation. In contrast, cells transformed with only pET plasmid show no aggregates.

Aggregation was also studied using a tube settling assay. Cultures of *E.coli*, transformed with either empty pET or HadA/na-containing pET, were incubated to late exponential phase and were then allowed to settle for 4 hours at room temperature. The HadA-expressing bacteria lost turbidity, but the control cells did not (Figure 35), indicating that HadA promotes bacterial aggregation.

5

10

Adhesion and invasion experiments were also performed with E.coli expressing HadA/na and a monolayer of Chang cells. Adherence (invasion) was calculated by counting the number of adherent (invaded) bacteria on cell monolayers (MOI = 1:1000). Results were taken as the mean \pm standard error of the mean of measurements made in triplicate, and are shown in Figure 29. The numbers of cells showing adhesion and invasion were as follows:

	Adherent	Invasive
HadA/na	1250 <u>+</u> 344 x 10 ⁴	26.3±6.6 x 10 ⁴
Empty plasmid	$10.5\pm2.1\times10^4$	$1.5\pm0.9 \times 10^4$

Adhesion and invasion were confirmed by immunofluorescence microscopy analysis (Figure 30).

Extracellular bacteria (green) and intracellular bacteria (red) can be seen.

Further studies of HadA include: construction of an isogenic HadA knockout of BPF Haemophilus influenzae strains for testing in adhesion/invasion assays; testing such knockout mutants to see if adhesion can be complemented by a NadA knockin; competition experiments with HadA and NadA in adhesion/invasion to human cells to see if HadA and NadA bind the same receptor.

It will be understood that the invention has been described by way of example only and modifications may be made whilst remaining within the scope and spirit of the invention.

REFERENCES (the contents of which are hereby incorporated by reference)

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